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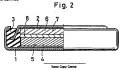
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- Non-aqueous secondary cell.

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NON-AQUEOUS SECONDARY CELL

Background of the Invention

(1) Field of the Invention

The present invention relates to a non-equeous secondary cell in which lithium or lithium alloy is used as the active material for the negative electrode, and particularly to an improvement in the positive electrode.

(2) Description of the Prior Art

Molybdenum trioxide, vanadium pentoxide and niobic sulfide have been proposed as the active material for the positive electrode of this type of secondary cell, but these substances have not been put to practical use to date.

For the positive electrode of the non-equeous primary cell, on the other hand, manganiese dioxide and carbon fluoride are income to be typical examples of active material and are actually employed for the purpose. Manganise dioxide has the advantages of being excellent in storage characteristics, abundant in the auth and inexponentive.

As the crystal structure of manganese clioside suited for the positive electrode, -y & MrOs health trained at temperatures of 250-050°C has been proposed as in Japanese Plantie Publication No. 40-25571. This y-\$MrOs, however, is unstalfactory in reversibility and has the problem of towering charge and discharge characteristics. The reason will be explained with reference to Figs. 1a-1e of the accompanying densings shrowly favering through X-reg (Iffract).

tion patterns. Fig. 1a shows a diffraction pattern prior to charging and discharging. Figs. 1b and 1c show diffraction patterns at a 10th discharge and charge, respectively. Compared with the pattern of Fig. 1a. it will be seen that the patterns of Figs. 1b and 1c show the angles of diffraction shifting to e lower side and the peaks becoming less sharp. These trends are more conspicuous and the peaks are almost leveled out in the patterns at a 100th discharge and charge shown in Flgs. 1d and 1e. respectively. It may be deduced from the above that a repetition of charges and discharges results in widening of the bond length between manganese and oxygen and in loosening of the crystal structure of manganese dioxide. Consequently, the manganese dioxide has poor reversibility and charge and discharge characteristics.

This applies also to β -Mn0₂ heat-treated at temperatures of 350-430°C as disclosed in U.S. Patent 4.133.856.

Thus, manganese dioxide is desirable as the active material for the positive electrode of the nonsqueous secondary cell, but involves difficulties in position.

Summary of the Invention

The object of the present invention, therefore, is to improve the reversibility of managenes closed without impairing the advantages of marganese ideode, i.e. excellent stronge characteristics, availability in abundance and low cost, thereby to improve charge and discharge cycle characteristics of the non-equeous manganese closide-lithium secondary cell.

The above object is fulfilled by a repeatedly chargeable and dischargeable non-aqueous secondary cell comprising a negative electrode having libitum or libitum alloy as an active material, a positive electrode having management discides as is active material, the possitive electrode including Libitum, as experient disposed between the positive electrode and the negative electrode, and e non-approximate forms of the property of

The above LLMnO₃ is obtained by heat-treating e-mixture of manganese dioxide and lithium salt. Specifically, the lithium salt is selected from the group consisting of lithium hydroxide, lithium nitrate, lithium phosphate, lithium carbonate and lithium oxide.

The lithium selt and manganese clioxide are mixed in a lithium-manganese moler ratio range of 10:90 to 70:30.

The mixture of manganese dioxide and lithium

salt is heat-treated in a **Iemperature range of 300-430°C, preferably 350-430°C.
The positive electrode is formed by heat-treating a mixture of manganese clicide and tithium salt to produce manganese clicide and LiMnOs, thereafter adding a conductive agent and a bindey, and

pressurizing and heal-treating a resulting mixture. The positive electrode may also be formed by heat-treating a mixture of manganese clication and lithium self to produce manganese clication and LighthOs, thereafter adding further manganese clication, a conductive agent and e binder, and pressurizing mixture and pressurizing mixture. Alternatively, the positive electrode mey be formed by heat-treating a mixture of manganese dicoide and lithium sait to produce LiMnO₂, therefiber adding manganese dicoide, a conductive agent and a binder, and pressurizing and heat-treating e resulting mixtures.

According to the present invantion, the positive electrode includes LIAMOS in addition to manganess dioxide constituting the active material. This composition prevents widening of the bond length between manganess and oxygen in the manganess dioxide and loosening of the crystal structure of the manganess addicted. Thus, the present invention provides an improvement in the reversibility and charges and discharge characteristics.

Brief Description of the Drawings

Figs. 1a-1e are views showing diffraction pattern where charges and discharge are repeated on a cell having a positive electrods formed of ~-8MnOs, Mroori Fig. 1a thowe a diffraction pattern prior to charging and discharging, Figs. 1b and 1c show diffraction pattern at 10th discharge and a subsequent charge, respectively, and Figs. 1d and 1e show diffraction patterns at a 100m discharge and a subsequent charge, respectively.

Fig. 2 is a half section of a cell according to the present invention, Fig. 3 is a view showing charge and di-

scharge characteristics of the cell.

Figs. 4a-4e are views showing diffraction petterns where charging and discharging are repeated on the cell according to the present invention, wherein Fig. 4e shows e diffraction pettern port to charging and discharging. Figs. 4b and 4c show diffraction petterns at a 10th discharge and a subsequent charge, respectively, and Figs. 4d and 4e show diffraction petterns at e 100th discharge and e subsequent charge, respectively.

Figs. 5 through 9 are views showing diffraction patterns for checking heat treatment temperature conditions for preparing LiAMnOs, where heat treatment is effected at temperatures of 250-C, 300°C, 375°C, 490°C, and 500°C, respectively.

Figs. 10 through 16 are views showing diffraction patterns for checking moler ratios between Li and Mn for producing LJsMnO_x where Li and Mn are mixed in the ratios of 5:95, 10:90, 20:80, 30:70, 50:50, 70:30, and 85:15, respectively.

Detailed Description of the Invention

EXAMPLE 1

A first example embodying the present invention will be described hereinafter with reference to a flet type non-aqueous socondary cell as shown in Fig. 2.

The Illustrated cell comprises positive and nec-

athve care 1 and 2 formed of stainless steel and separated from each other by an insulating packing 3 formed of polypropylens. Number 4 indicates a positive electrodic constituting the girls of the Invention, which is pressed upon e positive collector 5 essecuted to a bottom inside authea of the positive can 1. Number 6 indicates a negative electrodic pressed upon negative collector 7 secured to a pressed upon negative collector 7 secured to a pressed upon negative collector 7 secured to a formation of the positive collector of the positive ber 6 indicates a separator comprising a pressurmentative of polypropriser. This coll temploys as

electrolyte comprising lithium perchlorate dissolved in 1 molliter in e solvent mixture of propylene carbonate and dimethoxyethane.

The positive and negative electrodas are prepered as follows:

80 grame of chemical manganese dioxido having an average perticle size not exceeding 30 micron and 20 grams of situm hydroxide are first mixed in o morter, and then heat-treated in the air at 375°C for twenty hours. This heat treatment produces e mixture in which manganese dioxide

and Li₂MnO₃ coexist. The reaction formula for the Li₂MnO₃ production is as follows: MnO₂ + 2UOH -> Li₂MnO₃ + H₂O (1)

Next, the active material powder thus obtained in mixed with acetylene black acting as conductive agent and fluoric resin powder acting as binder in a weight ratio of 90.5% to produce a bland for forming the possible electrods. This bland is moisted under a pressure of 2 transcribt into a shape having a 20mm dismotor, and then health-resided at 200°C, whereby the possible electrods is completed. The necessity electrods on the other head is promoted.

by punching e piece 20mm in diameter out of e tithium foil having a selected thickness. Incidentally, the cell le 24.0mm in diameter and 3.0mm in thickness. This cell embodying the present invention is hereinafter referred to as Call

A1.

A positive electrode is produced in the same way as in EXAMPLE 1 except that 60 grams of lithium carbonate are used instead of the 20 grams of lithium hydroxide. This cell is hereingther re-

ferred to as Coll A2

The reaction formula for the Li₂MnO₃ production in this example is as follows: MnO₃ + Li₂CO₃ → Li₂MnO₃ + CO₂

COMPARATIVE EXAMPLE 1

A positive electrode is produced in the same way as in EXAMPLE 1 except that the lithium salt is not added. This cell produced for comparison purposes is browingfair referred to as Cell B1.

COMPARATIVE EXAMPLE 2

Manganese dioxide is added and mixed with lithium hydroxide as in EXAMPLE 1 but, as distinct from EXAMPLE 1, the mixture is not heat-treated here. The coil thus produced is hereinsfiler referred to ac Call Rs.

Fig. 3 shows charge and discharge cycle characteristics of these cells. The data were obtained from the conditions that the discharge was carried out in a current of 3mk for four hours, the charge in the current of 3mA, and the charge ending voltage was 4.0°V.

It will be seen from Fig. 3 that discharge ending vollages of Coles B1 and B2 drop champly around the 100th cycla whereas those of Coles A1 above sharp drops up to the Viroling of 100 Coles above sharp drops up to the Viroling of 100 Coles above sharp drops up to the Viroling of 100 Coles Americansistics. Firm the characteristics of Cells B1 and B2 produced for comparison purposes, it is understood that the addition of Billiams salt would not produce a satisfactory result for Improvise of Viroline Cells B1.

The reason for the improvement in the cycle characteristics will be explained next with reference to Figs. 4a-4e.

Compand with the a diffraction pattern prior of charging and discharge above in Fig. 4, diffraction patterns at a 10th cycle discharge and a sixtem of the companion of a 10th cycle discharge and a solitoncian argument of the companion of the companion of strates that the load length between marganisms and congen does not increase and the crystal strates that the load length between marganisms and congen does not increase and the crystal strates that the load length between marganisms and congenity of the companion of the companion strates and the companion of the companio Heat treatment temperatures for producing LightCo, have been checked, and the results will be described hereinafter referring to the X-ray diffraction patterns shown in Figs. 5 strough 9. In threes tests lifthium and manganess were mixed in a fixed ratio of 30:70, and LiOH was used as the lithium saft and MnO. as the manganese oxide.

When the heat treatment was carried out at 20°C, Light's, was not produced at all as shown in Figs. 5. The reason is considered that no reaction stops place between LOH and MoV, at that temperature is inadequate for the purpose of the temperature is inadequate for the purpose of the present invention. When the heat treatment was carried out at 300°C, 57°C and 45°C, Light's, or special control of the property of

and MnO₅, and only MinO₇ remains after all LOH has reacted with MnO₅. Consequently, the hast treatment in the temperature range of 300-450°C produces the effect of the invention to the full. Further, when the heat treatment was certed out at 500°C, LSMnO₅ we sproduced and MnO₇ meminds, but MnO₅ which is an undestrable cell material was produced at a result of decomposition of MnO₅ as shown in Fig. 9. Consequently the heat treatment at this temperature fails to produce the effect of the

invention.

The above test results prove that the desirable temperature range for the heat treatment is from 300°C in 430°C.

The heat treatment carried out in the temperature range of 300-430°C has the advantage of producing LisMnO₃ and dehydrating MnO₂ acting as the active material.

Where the hest treatment for producing LiMnO₃ is simed at removal of combined water in manganese dioxide, it is desirable to carry out the hest treatment in the temperature range of 350-

Furthermore, malar ratios between fishum and marganese for producing LiMnOs, were chacked and the results will be described hereinathor referring to the X-ray diffraction patients shown in Figs. 10 innough 16. In these test the heat treatment temperature was fend to 375°C, and LIOH was used as the tithium sait and MnOs as the manganese oxide. When tithium sait and MnOs as the manganese were mixed in a

moler ratio of 5:95, LishinCo, was not produced at all as shown in Fig. 10. Consequently, the moler ratio of 5:95 between lithium and manganese does not accomplish the purpose of the present invention. When the lithium and manganese were mixed in moler ratios of 10:90, 20:90, 30:70, 50:50, and 70:30, LishinCo, was produced as shown in Figs. 11 hrough I.S. responsively. This is because, in this range of mixing rotice, the reaction supposed in the formule (1) takes place between LOH and McC. Consequently, mixing of thism and manufactures of the control of th

The above test results prove that the desirable molar ratio range between Ethium and manganese is from 10:90 to 70:30.

When Ethium and mangenese were mixed in the motar ratio of 703.0 Lishfox) was produced but MnO₂ was not as shown in Fig. 15. Even so, the purpose of the invention will be stiffled by a clied MnO₂ after producing Lishfox, Further, it is possible to vary the ratio between Lishfox of and MnO₂ by mixing lithium and manganese in the molar ratio case of 10.86 by 70.30 to produce Lishfox, and MnO₂ and

thereafter adding MnOs.
In obtaining LieMnOs by heat-treating the mixture of manganese dioxide and lithium salt as in this invention, the lithium salt is not limited to those given in the foregoing examples but may comprise

lithium nitrate, lithium phosphate or lithium oxide.

The type of manganase dioxide is not limited to chemical manganese dioxide but may of course comprise natural manganese dioxide or electrolytic

mangenese dioxide.

Furthermore, the present invention is applicable not only to the non-aqueous electrolyte cell but to the solid electrolyte cell also.

Claims

- A repeatedly chargeable and dischargeable non-acusous secondary cell comprising:
- e negative electrode having lithium or lithium alloy as an active material, e positive electrode having manganese dioxide
- e possive electrode naving manganese dioxide as an active material, said positive electrode including Li₂MnO₃.

 a secarator disposed between said positive
- electrode and said negetive electrode, and a non-aqueous electrolyte.
- A cell as claimed in claim 1 wherein said Li₃MnO₂ is obtained by heat-treating a mixture of manoanese dioxide and lithium salt.
- A celt as claimed in claim 2 wherein said lithium selt is selected from the group consisting of lithium hydroxide, lithium nitrate, lithium phosphate, lithium carbonate and lithium oxide.

- A cell as claimed in claim 2 wherein said lithium sall and manganese dioxide are mixed in a lithium-manganese molar ratio ranga of 10:90 to 70:30.
- A cell as claimed in claim 2 wherein said mixture of manganese dicoide and lithium sait is heat-treated in a temperature range of 300-430°C, preferably 350-430°C.
- 6. A cell as claimed in claim 1 wherein said positive electrode is formed by heat-treating e mixture of manganese dioxide and lithium sait to produce manganese dioxide and LisMnOs, thereafter adding e conductive agent and a binder, and pressurizing and heat-treating a resulting mixed.
- 7. A coil as claimed in claim 1 wherein said positive electrode is formed by heat-treating a mixture of manganess dioxide and tithium sait to produce manganess dioxide and LishnOs, thereafter adding-further manganess dioxide, a conductive soort and a binder, and pressurting and heatt-part of the conductive of the conductive conductive.
- treating a resulting mixture.

 8. A cell as claimed in claim 1 wherein said positive electrode is formed by heat-treating a mixture of manganese cloxide and lithium saft to produce LijkinO₃, thereafter adding manganese diox
- ide, a conductive agent and a binder, and pressurizing and heat-treating a resulting mixture. 9. A cell as citimed in claim 1 wherein said
- negative electrode is selected from the group consisting of pure lithium, lithium-aluminum alloy and lithium-magnesium alloy. 10. A cell as claimed in claim 1 wherein said separator comprises a porous membrane of poly-
- propylene. s 11. A cell as claimed in claim 1 wherein said alactrolyte comprises e liquid mixture formed by dissolving lithium perchlorate in e solvent mixture
- of propylene carbonate and dimethoxyethane.

 12. A cell as claimed in claim 1 wherein said opositive electrosis to pressed upon a positive collector secured to a bottom inside surface of a costitive terminal can.
- 13. A cell as claimed in claim 1 wherein said negative electrode is pressed upon a negative collector secured to a bottom inside surface of a negative terminal can.
 - negarre termins can.

 14. A repeatedly chargeable and dischargeable
 non-aqueous secondary cell comprising;
 e negative electrode having lithium or lithium
 - alloy as an activa material, e positive electrode having a mixture in which manganese dixxide as an active material and
- a separator disposed between said positiva se electrode and said negative electrode, and e non-aqueous electrolyte.

15. A cell as claimed in claim 14 wherein said positive electrode is formed by heat-treating e mixture of manganese dloxide and lithium salt to produce manganese dioxide and LisMnOs, thereafter adding a conductive egent and a binder, and pres-

surizing and heat-treating a resulting mixture. 16. A cell as claimed in claim 14 wherein said positive electrode is formed by heat-treating a mixture of manganese dioxide and Ethium salt to produce Li₂MnO₃, thereafter adding manganese dioxide, a conductive agent and a binder, and pres-

surizing and heat-treating a resulting mixture. 17. A cell as claimed in claim 14 wherein said negative electrode is selected from the group consisting of pure lithium, lithium-aluminum alloy and lithium-magnesium alloy.

18. A cell as claimed in claim 14 wherein said separator comprises a porous membrane of poly-

propylene. 19. A cell as claimed in claim 14 wherein said electrolyte comprises a liquid mixture formed by dissolving lithium perchlorate in a solvent mixture of propylene carbonate and dimethoxyethene.

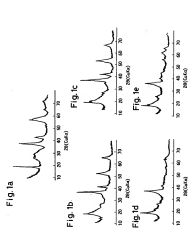
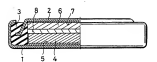
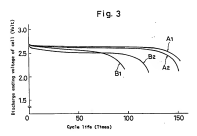
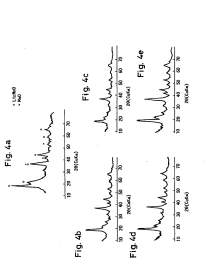


Fig. 2

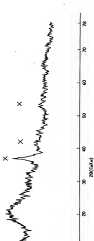


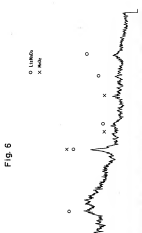




O L12Mn03 X Mn02

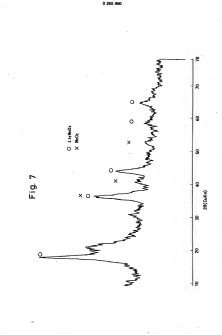
Fig. 5

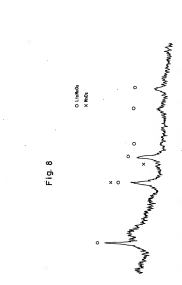




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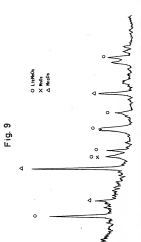
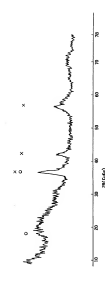
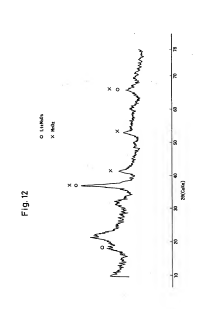




Fig. 10







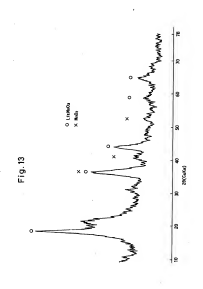
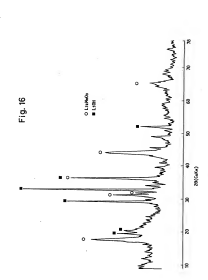


Fig. 14

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DOCUMENTS CONSIDERED TO BE RELEVANT				EP 87115912.5
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A	GMELINS HANDBUC CHEMIE, 8th edi 56, "Mangan, Te dungen", 1975	H DER ANORGANISCHEN tion, System-no. il C2, Verbin-	1-8,15, 16	
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-				TECHNICAL FIELDS SEARCHED (IN. CI.4)
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	The present search report has b	een drawn up for all claims		

CATEGORY OF CITED DOCUMENTS